IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of Hajime KONDO

Application No.: 10/558,389

Filed: November 29, 2005

For: MODIFIED NATURAL RUBBER OR MODIFIED NATURAL RUBBER LATEX, AND

RUBBER COMPOSITION AND PNEUMATIC TIRE

Group Art Unit: 1796

Examiner: Peter D. Mulcahy Confirmation No.: 5714

DECLARATION UNDER 37 C.F.R. § 1.132

I, Makiko Yonemoto, declare that:

I am a co-worker of Mr. Hajime Kondo who is the inventor of the above-captioned patent application.

I received my Master of Science and Engineering from Tokyo Institute of Technology in 2004, and have been employed by Bridgestone Corporation since 2004, where I have been engaged mainly in research and development of new natural rubber.

I have made the following experiments in order to evaluate the Mooney viscosity, the ultimate strength (fracture resistance), the crack growth resistance and the tan δ of the rubber composition comprising the modified natural rubber, which is obtained by graft-polymerizing natural rubber latex with a polar group-containing monomer and then coagulating and drying, and carbon black and/or silica, wherein a grafting ratio of the polar group-containing monomer is 0.01-5.0% by mass per the natural rubber latex, and a content of the modified natural rubber in a rubber component of the rubber composition is at least 15% by mass. Further, I have made the following experiments in order to evaluate the rolling resistance of the tire using the above rubber composition.

Experimental Procedure

<Production Example of Modified natural rubber A-1>

A modified natural rubber A-1 is obtained in the same manner as in Production Example 1 described in the present specification. As the modified natural rubber A-1 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this

example, 3.0 g of N,N-diethylaminoethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 3.0/600 = 0.5%.

<Production Example of Modified natural rubber A-2>

A modified natural rubber A-2 is obtained in the same manner as in Production Example 1 described in the present specification except that 28.2 g of N,N-diethylaminoethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber A-2 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 28.2 g of N,N-diethylaminoethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 28.2/600 = 4.7%.

<Production Example of Modified natural rubber A-3>

A modified natural rubber A-3 is obtained in the same manner as in Production Example 1 described in the present specification except that 34.8 g of N,N-diethylaminoethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber A-3 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 34.8 g of N,N-diethylaminoethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 34.8/600 = 5.8%.

<Production Example of Modified natural rubber A-4>

A modified natural rubber A-4 is obtained in the same manner as in Production Example 1 described in the present specification except that 42.0 g of N,N-diethylaminoethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber A-4 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 42.0 g of N,N-diethylaminoethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 42.0/600 = 7.0%.

<Pre><Pre>roduction Example of Modified natural rubber B-1>

A modified natural rubber B-1 is obtained in the same manner as in Production Example 1 described in the present specification except that 3.0 g of 2-hydroxyethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber B-1 is analyzed in the same manner as in Production Example 1 described in the present specification,

it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 3.0 g of 2-hydroxyethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 3.0/600 = 0.5%.

<Production Example of Modified natural rubber B-2>

A modified natural rubber B-2 is obtained in the same manner as in Production Example 1 described in the present specification except that 26.4 g of 2-hydroxyethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber B-2 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 26.4 g of 2-hydroxyethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 26.4/600 = 4.4%.

<Production Example of Modified natural rubber B-3>

A modified natural rubber B-3 is obtained in the same manner as in Production Example 1 described in the present specification except that 31.2 g of 2-hydroxyethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber B-3 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 31.2 g of 2-hydroxyethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 31.2/600 = 5.2%.

<Pre><Pre>roduction Example of Modified natural rubber B-4>

A modified natural rubber B-4 is obtained in the same manner as in Production Example 1 described in the present specification except that 42.0 g of 2-hydroxyethyl methacrylate is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber B-4 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 42.0 g of 2-hydroxyethyl methacrylate is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 42.0/600 = 7.0%.

<Production Example of Modified natural rubber F-1>

A modified natural rubber F-1 is obtained in the same manner as in Production Example 1 described in the present specification except that 3.0 g of acrylonitrile is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber F-1 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 3.0 g of acrylonitrile is added to 600 g of latex (60% of 1000 g of the concentrated

latex), so that the grafting ratio is 3.0/600 = 0.5%.

<Production Example of Modified natural rubber F-2>

A modified natural rubber F-2 is obtained in the same manner as in Production Example 1 described in the present specification except that 27.0 g of acrylonitrile is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber F-2 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 27.0 g of acrylonitrile is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 27.0/600 = 4.5%.

<Production Example of Modified natural rubber F-3>

A modified natural rubber F-3 is obtained in the same manner as in Production Example 1 described in the present specification except that 33.0 g of acrylonitrile is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber F-3 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 33.0 g of acrylonitrile is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 33.0/600 = 5.5%.

<Production Example of Modified natural rubber F-4>

A modified natural rubber F-4 is obtained in the same manner as in Production Example 1 described in the present specification except that 42.0 g of acrylonitrile is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber F-4 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 42.0 g of acrylonitrile is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 42.0/600 = 7.0%.

<Production Example of Modified natural rubber O-1>

A modified natural rubber O-1 is obtained in the same manner as in Production Example 1 described in the present specification except that 0.6 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-1 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 0.6 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 0.6/600 = 0.1%.

<Production Example of Modified natural rubber O-2>

A modified natural rubber O-2 is obtained in the same manner as in Production Example 1 described in the present specification except that 3.0 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-2 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 3.0 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 3.0/600 = 0.5%.

<Production Example of Modified natural rubber O-3>

A modified natural rubber O-3 is obtained in the same manner as in Production Example 1 described in the present specification except that 28.8 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-3 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 28.8 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 28.8/600 = 4.8%.

<Production Example of Modified natural rubber O-4>

A modified natural rubber O-4 is obtained in the same manner as in Production Example 1 described in the present specification except that 60.0 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-4 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 60.0 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 60.0/600 = 10%.

<Production Example of Modified natural rubber O-5>

A modified natural rubber O-5 is obtained in the same manner as in Production Example 1 described in the present specification except that 90.0 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-5 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 90.0 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 90.0/600 = 15%.

<Production Example of Modified natural rubber O-6>

A modified natural rubber O-6 is obtained in the same manner as in Production Example

1 described in the present specification except that 33.0 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber Q-6 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 33.0 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 33.0/600 = 5.5%.

<Production Example of Modified natural rubber O-7>

A modified natural rubber O-7 is obtained in the same manner as in Production Example 1 described in the present specification except that 42.0 g of 2-vinylpyridine is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-7 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 42.0 g of 2-vinylpyridine is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 42.0/600 = 7.0%.

<Production Example of Modified natural rubber U-1>

A modified natural rubber U-1 is obtained in the same manner as in Production Example 1 described in the present specification except that 126.0 g of γ -methacryloxypropyl trimethoxy silane is used instead of 3.0 g of N,N-diethylaminoethyl methacrylate. As the modified natural rubber O-1 is analyzed in the same manner as in Production Example 1 described in the present specification, it is confirmed that 100% of the monomer added is introduced into the natural rubber molecule. In this example, 126.0 g of γ -methacryloxypropyl trimethoxy silane is added to 600 g of latex (60% of 1000 g of the concentrated latex), so that the grafting ratio is 126.0/600 = 21%.

<Pre><Pre>reparation of Rubber composition>

A rubber composition is prepared by using the modified natural rubber A-1, A-2, A-3, A-4, B-1, B-2, B-3, B-4, F-1, F-2, F-3, F-4, O-1, O-2, O-3, O-4, O-5, O-6, O-7 or U-1, respectively, according to a compounding recipe shown in the following Tables I, II, III, IV, V and VI. With respect to the resulting rubber composition, the Mooney viscosity, the ultimate strength (fracture resistance), the crack growth resistance and the tan δ are evaluated according to the following methods. Further, with respect to a tire prepared by using the rubber composition, the rolling resistance is evaluated according to the following method. Results are shown in Tables I, II, III, IV, V and VI.

(1) Mooney viscosity

For evaluating the processability of the rubber compositions, unvulcanized rubber

samples were kneaded together with a vulcanized ingredient and preheated at 130°C for 1 minute using "MOONY VISCOMETER SMV201" manufactured by Shimadzu Co., Ltd., and then the rotation of the rotor was started, and the value after a lapse of 4 minute was measured as ML₁₊₄. The higher the value, the higher the unvulcanized viscosity and lower the processability or productivity.

(2) Ultimate strength (fracture resistance)

A specimen vulcanized at 160°C for 20 minutes was measured in accordance with JIS K6251-1993 to determine the tensile strength at 23°C. The higher the value, the higher the fracture resistance.

(3) Crack growth resistance

Dumbbell-shaped specimens were punched out of a rubber composition vulcanized under vulcanization conditions at 160°C for 10 minutes, and the specimens having formed a preliminary crack of 5 mm at the center thereof were mounted on a fatigue testing machine, and subjected to strokes at 7 Hz at a constant stress, a temperature of 80°C, and a chuck distance of 20 mm, and the number of cycles necessary to a complete fracture was represented as an index on the basis that the case of Additional Comparative Example 5 is 100. The higher the value, the longer the life and the better the crack growth resistance.

(4) tan δ

Measurements were conducted on a rubber composition vulcanized under vulcanization conditions at 160°C for 14 minutes using a spectrometer (dynamic viscoelasticity measuring testing machine), at an initial load of 160 g, a frequency of 50 Hz, a strain of 1%, and a measuring temperature of 23°C. The value was represented as an index on the basis that the case of Additional Comparative Example 5 is 100. The larger the index value, the lower the loss.

(5) Rolling resistance

A rubber composition was used as the ply coating rubber, and the rolling resistance of the tire (size: 185/70R14) was measured with a drum, and the value was represented as an index on the basis that the case of Additional Comparative Example 5 is 100. The higher the value, the better the rolling resistance and the better the low rolling resistance.

		FI	Table I			7	A 3.724
			Additional	Additional	Additional	Additional Ormanative.	Additional
			Example 1	8	9	Example 11	Example 12
	Natural rubber (grafting ratio=0%)		100	•	,	J	•
	Modified natural subbot A-1 (grafting		ı	100	ı	9	ı
	Modified natural nubber A-2 (graffing		1		100	ı	1
	ratio=4,7%)						
<u></u>	Modified natural rubber A-3 (graffing		•	•	,	100	•
	ratio=5.8%)	Darts					
	Modified natural nibber A-4 (grafting	<u></u> , 25	1	•	ì	ı	8
	ratio=7.0%)	38E				{	£
	Carbon Black	TOTAL TOTAL	R	SS	3	7	7
	CHS.			L	1		1
			12	77	23	77	12
	Spiriting of the state of the s		15	15	15	15	15
	Searchtau 20 0 11		4	4	4	4	4
	ZITC CAME		0.8	0.8	0.8	89	0.8
	VIKAIIZAIIJITAMAAAA		cr.	m	m	3	ന
	Summ		Q	82	7%	ξ.	83
	Mooney viscosity	.	3 6	8	7	E	18
	Ulfimate strength	MITA.	30,	3 8	ξ	15	6
	Crack growth resistance		3	3	777	707	Live P
	121 S	index	105	25	248	25	/51
	Dollivanscidance	index	81	105	106	102	102
•	KOMPETENSIAN		- 4.5	1	13 con	in the second series	ith

* Modified natural rubbers A-1, A-2, A-3 and A-4 are a modified natural rubber graff-polymerized with N,N-diethylaminoethyl methacrylate.

-		띰	Table II				
			Additional	Additional Additional	Additional	Additional	Additional
		,	Comparative Fearmole 1	Example 10	Example 11	Comparative Example 13	Comparative Example 14
	(J00 - 2 - 1 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2	1	100			,	,
	Natural nubber (gramme rank—170)		3				
	Modified returnitubber B-1 (grafting	•	•	83	r)	•
	Identify (1.1 m of the control of t						
	Modified natural rubber B-2 (gramug		1	ı	8	ı	ì
	ratio=4.4%)						
	Modified natural reboer B-3 (graffing		1	ı	,	100	1
	ratio=5.2%)	1					
,	Modified natural nabber B4 (graffing		,	,	, 1	1	100
Formulation	ratio=7.0%)	5					8
	Carton Hark	SP	ଝ	30	ଷ	R	3
	Calcal		1	8	ଳ	8	ଚ
	Suita Si Suita Suita Suita Suita Suita Suita Suita Suita Suita Suita Suita Suita Suita Si Suita Suita Si Si Suita Si Si Si Si Si Si Si Si Si Si Si Si Si	-L-^-	12	12	12	77	17
	Single Cil	~~~	7	15	15	15	15
	SteancAcid	******		1	T	4	4
	ZincOxide		1 00	- 02	80	08	0.8
	Volcanization Accelerator		97	OM)	3	6	c
	C. Her		m	co.	3	'n	า
	N. A. Constant Control of the		83	88	æ	88	&
	INICIES VISCOLD	2	%	25	23	13	17
	Ultimate strength	יין די	3 2	پې	101	8	35
Results	Crack growth resistance	Yani.		3 6	ž	148	147
	tanô	H	COT	3	3	2 2	20,
	Dollmanschance	index	. 001	8	107	<u>a</u>	<u>a</u>
	Name of the second of the seco			1		- a - alymonized with	4

* Modified natural rubbers B-1, B-2, B-3 and B-4 are a modified natural rubber graft-polymerized with 2-hydroxyethyl methacrylate.

			Aridificanal	Additional	Additional Additional	Additional	Additional
			Commentive	Example	Example	Comparative	Comparative
			Example 1	12,	ŧΞ	Example 15	Example 16
	Natural nibber (graffing ratio=0%)		100		Ŀ	3	
	Motified ratural nubber F-1 (graffing			100	1	ı	1
	xafio=(1.5%)						
	Modified natural rubber F-2 (grafting		ſ	1	130	ì	ı
	rafio=4.5%)						
	Modified natural rubber F-3 (grafting		•	1	1	100	
	ratio=5.5%)	1					
•	Modified natural rubber F4 (grafting	<u>4</u> , j		,	,	ı	100
Formulation	ratio=7,0%)	<u> </u>					
	Carton Black	STEEL STEEL	ନ	R	ଷ	R	83
	Cilica		1	æ	R	30	ଝ
	Sanda Off	-	12	23	²³	12	12
	Charles Ariel	γ	15	15	21	1.5	15
	Tra Ocida	.	4	4	च	Þ	4
	Videnmetica Amelegia	7	80	89	0.8	0.8	89
	Suffin		m	ξŲ	3	es	3
	Momentaismeits		98	ळ	88	88	83
	Titionale channelly	25	38	75	22	18	17
1 1	Charle sucretty recipients	index	105	83	101	85	፠
SINCE	ten S	index	33	148	153	147	145
	Dollarovitona	index	100	392	106	· 105	105

* Modified natural rubbers F-1, F-2, F-3 and F-4 are a modified natur

Table IV

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			Additional	Additional	Additional	Additional	Additional	Additional	Additional
			Comparative	Example	Example	Example	Example	Comparative	Comparative
			Example 1	1	2	'n	4	Example 17	Example 18
	Natural milyber (graffing ratio=0%)		100	,	•	ı	,	•	*
	Modified natural rubber O-1 (grafting	.	1	100	ı	ı	•	ı	ı
	ratio=0.1%)								
	Modified patural rubber 0-2 (grafting		r	t	901	•	1	ı	1
	ratio=0.5%)								
	Modified natural rubber O-3 (graffing		ı		ı	100	100	ı	1
	ratio=4.8%)								· ·
	Modified natural nubber 0-6 (grafting	show	1	,	ı	ſ	ì	9	ı
	zafo=55%)								
Formulation	Formulation Modified natural nabber 0-7 (grafting	LOY TIESS	,	1	1	ı		1	100
	ratio=7.0%)						1	•	8
	Carbon Black		SS.	ଟ	83	8	3	ਨ	7
	Silin		1	1	1	ଝ	•		
	الماسية	***	12	12	12	12	12	12	12
	Species Activity		15	31	15	15	15	1.5	15
	Zan Onida		4	4	4	4	7	4	₩
	Tiloningin Avalentin	••	80	0.8	83	83	80	0.8	33
	Vulcancia Curaciano.		3	ť	3	3	3	3	3
	Montestations		89	ج	88	23	74	IJ	86
	The strength	Z.	38	25	27	83	24	21	19
7	Out month resistance	1	3	100	103	108	102	101	86
	the S	ig.	193	125	140	155	146	138	137
	Dollhomedstance	işi.	100	101	165	107	105	102	102
	Number of the second se							•	

* Modified natural rubbers O-1, O-2, O-3, O-6 and O-7 are a modified natural rubber graft-polymerized with 2-vinylpyridine.

Table V

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			Additional	Additional	Additional	Additional
			Comparative	Comparative	Comparative	Comparative
			Example 1	Example 2	Example 3	Example 4
	Natural rubber (grafting ratio=0%)		100		•	
	Modified natural nubber 04 (graffing	,		100	3	,
	ratio=10%)					
	Modified natural nubber 0-5 (grafting		ı		100	ı
	ratio=15%)					
	Modified natural nubber U-1 (grafting	rante.	1	,	•	100
1	ratio=21%)	3, 4				
Formulation	Carism Black	'n	SS.	50	RS.	R
		THESS	1	1	•	33
	Chindle ()	,	27	12	12	12
	Creary Arid	•	51	15	15	1.5
	Zur Oxide		4	ŧ	4	4
	Visionization Accelerator		870	80	0.8	038 0
	Criffy		'n	m	3	.3
	Momentalenette		69	81	\$	25
	I Himste Strength	MPa	25	18	17	16
Domite	Crack omworth resistance	index	105	25	83	93
	tanô	index	105	135	130	150
	Relimensistance	index	100	103	103	걸
			1. C. J. L. J. L.	Las Amely	transperiment and	٠.

* Modified natural rubbers O-4 and O-4 are a modified natural rubber graft-polymerized with

2-vinylpyridine.

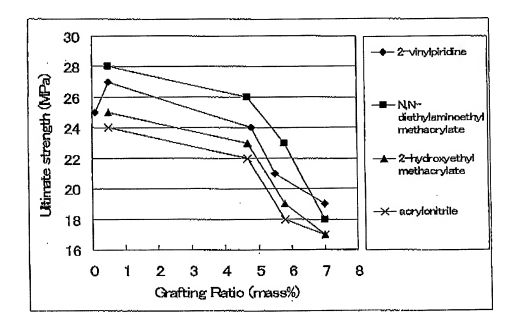
* Modified natural nubber U-1 is a modified natural rubber graft-polymerized with γ -methacryloxypropyl trimethoxy silane.

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<u>VI</u>	Additional Additional Additional Additional Additional	Comparative Example Comparative Example Omparative Comparative Example 7 Example 10	40 - 20 - 10 -	. 40 . 20 . 10	60 68 80 90 90	50 50 50 50 50	1	12 12 12 12 12	15 15 15 15 15 15	4 4 4 4 4	0.8 0.8 0.8 0.8 0.8	3 3	56 55 56 54 54	21 21 20 18 17	95 92 93 91 90	95 110 85 90 85 85	99 101 96 99 97 97	
Table VI	Additional	Example 5	•	70	8	S.	ŧ	12	15	4	88	3	53	22	100	135	103	
	Additional	Omparative Example 6	202	•	88	20	30	12	15	4	80	3	23	83	,38	102	100	
	Additional	Comparative Example 5	Æ	,	93	8	,	12	15	4	80	m	8	25	100	100	100	
		~~~		T		State.	È,	Sem -						Z P	index	À L	ig.	
			Natural nubber (grafting ratio=1%)	Modified natural rubber	Charachatatiere mitter	Carbon Black	Silcs.	Spingle Oil	Siemic Acid	Zinc Oride	Villeanization Accelerator	Suffin	Moraecroismeilo	1 Throate chenoth	Contractify resistance	ton &	Rolling resistance	-
						•	Formulation					•••			Domition	CIIII CONT		

* Modified natural rubber O-2 is a modified natural rubber graft-polymerized with 2-vinylpyridine.

Further, the results of the ultimate strength of the Additional Examples 8 and 9 and the Additional Comparative Examples 11 and 12 in Table I, the results of the ultimate strength of the Additional Examples 10 and 11 and the Additional Comparative Examples 13 and 14 in Table II, the results of the ultimate strength of the Additional Examples 12 and 13 and the Additional Comparative Examples 15 and 16 in Table III, and the results of the ultimate strength of the Additional Examples 1, 2 and 4 and the Additional Comparative Examples 17 and 18 in Table IV are summarized in the following graph, respectively.



### (Summary)

As seen from Tables I, II, III and IV and the graph, when the grafting ratio of the modified natural rubber exceeds 5% by mass, the ultimate strength (fracture resistance) is largely deteriorated. Further, as from Tables I, II, III and IV, these unexpected results can be obtained in the tests using various modified natural rubbers.

I declare further that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with knowledge that willful false statements

and the like so made are punishable by fine or imprisonment, or both, under § 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Date: 2010, 4. 2 Declarant: Makiko Yonemoto

Makiko Yonemoto